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# A Bayesian Translational Framework for Knowledge Propagation, Discovery, and Integration Under Specific Contexts

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## Abstract

*The immense corpus of biomedical literature existing today poses challenges in information search and integration. Many links between pieces of knowledge occur or are significant only under certain contexts—rather than under the entire corpus. This study proposes using networks of ontology concepts, linked based on their co-occurrences in annotations of abstracts of biomedical literature and descriptions of experiments, to draw conclusions based on context-specific queries and to better integrate existing knowledge. In particular, a Bayesian network framework is constructed to allow for the linking of related terms from two biomedical ontologies under the queried context concept. Edges in such a Bayesian network allow associations between biomedical concepts to be quantified and inference to be made about the existence of some concepts given prior information about others. This approach could potentially be a powerful inferential tool for context-specific queries, applicable to ontologies in other fields as well.*

## 1 Introduction and Objective

The millions of published works of biomedical literature cover an enormous array of knowledge. Over 21 million articles are indexed in PubMed alone, and around 700,000 new articles are added yearly<sup>1</sup>. Additionally, data from millions of experiments are archived in diverse databases. The large size of today’s body of biomedical knowledge and swiftness with which new information is being added present challenges in organization and navigation. The rise of such a large amount of information in recent years is changing the nature of biological knowledge from a descriptive practice to a more data-driven one, and finding specific information through manual search is growing increasingly difficult.

Biomedical ontologies can potentially be used to address these challenges. Tremendous efforts have been made to create diverse ontologies that together include all biomedical concepts. The National Center of Biomedical Ontology (NCBO) BioPortal<sup>2,3</sup> provides over 250 such ontologies with over 5 million concepts<sup>6</sup> to researchers. Moreover, researchers use ontology terms to annotate experimental data and works of literature. Hence, an automated, efficient framework that navigates and integrates the information embedded in these ontological links would be a powerful research tool that utilizes an immense range of biomedical knowledge. However, ontologies are usually developed in a silo, and the separateness of ontologies has so far hindered the practical application of ontological organization. Hence, a crucial question remains unanswered: is it possible to automatically and efficiently use biomedical ontologies to infer new knowledge?

This work presents such an automated framework that integrates biomedical ontologies and infers knowledge from abstracts of literature and descriptions of experimental data in response to a user-defined query. In particular, this framework infers information particular to a given *context*, or situation. Context-specificity is useful because researchers often have questions relevant to specific situations, and the same biological concepts may be linked in some contexts but not in others. For example, two traits might not generally be observed together, but in the context of a specific genetic condition, they may coexist frequently. The proposed framework identifies these types of linkages.

## 2 Mapping Ontologies: An Overview

A logical first step is to integrate the disparate biomedical ontologies. We seek a reliable framework for mapping ontological relationships that (1) considers diverse types of relationships between terms, (2) accounts for uncertainty in ontology integration, (3) is scalable to the size of biomedical ontologies, and (4) is able to be tailored to specific contexts. So far, no such framework has been developed.

The Unified Medical Language System (UMLS)<sup>4</sup> has integrated over 2 million names for approximately 900,000 biological concepts. However, mappings of UMLS concepts were manually curated, so there remain inconsistencies and errors in the mappings, and it is difficult for mappings to keep pace with the rate at which knowledge is expanding<sup>5</sup>. Many non-manual methodologies exist for ontological integration, including semi-automatic methods such as PROMPT<sup>7</sup> and GLUE<sup>8</sup>, and

automatic methods such as IF-MAP<sup>9</sup>, ANCHOR-PROMPT<sup>10</sup>, and MAFRA<sup>11</sup>. In Chua et al.<sup>12</sup>, more than 30 ontology mapping methods are surveyed and categorized into 7 categories. However, almost all proposed methods are not publicly available or are not scalable to the size of biomedical ontologies<sup>13</sup>.

Two recent methods, Association Rule Ontology Matching Approach (AROMA)<sup>14</sup> and Lexical OWL Ontology Matcher (LOOM)<sup>13</sup>, are publicly available and easily scalable. These two methods differ significantly. LOOM is used for discovering equivalence correspondences between concepts, is based on lexical matching, and does not require text corpora to work. In contrast, AROMA is used for inferring subsumption relationships between concepts, is based on a statistical measure known as implication intensity, and requires additional text corpora. Though these methods are steps forward, they do not consider the inevitable uncertainty of ontology mapping.

Some ontology-mapping studies do consider uncertainty by incorporating probabilistic uncertainty into their description logic by using Bayesian networks<sup>15-19</sup>. For example, a framework called OMEN<sup>16</sup> creates Bayesian networks of ontologies by drawing initial probabilities from *a priori* knowledge and then using a set of meta-rules to determine conditional probabilities between nodes. The conditional probabilities represent influences induced by nodes on their children. Two other algorithms, MSBN<sup>17</sup> and AEBN<sup>18</sup>, create pairwise correspondences between semantically identical concepts and propagate information through these correspondences between two ontology-specific Bayesian networks. The algorithm BayesOWL<sup>15</sup> uses a process similar to those of MSBN and AEBN but is more comprehensive: it links similar concepts as well as identical concepts by defining the similarity of concepts probabilistically by their joint distribution. More methods for probabilistic modeling of uncertainty in linking ontologies can be found in Lukasiewicz<sup>20</sup>.

### 3 Context-Specific Ontology Mapping and Probabilistic Inference: A Novel Technique

The method presented in this paper is distinct from the aforementioned ontology-mapping methods in its use of a context-sensitive algorithm. In prior work, a context-specific mapping algorithm based on the Bayes factor<sup>21</sup> was developed. This study adapts and applies that mapping method to construct the backbones of context-centered Bayesian networks for inference about biomedical relationships. This context-specific mapping approach has three main advantages: (1) mappings created are specific to the question under investigation, so unrelated concepts are pruned; (2) inference is less prone to noise generated from considering many unrelated concepts and can be more accurate; and (3) pruning many irrelevant concepts allows the inference algorithm to be scaled to the large size of most biomedical ontologies.

Once the Bayesian backbone is constructed, probabilistic inference on the framework accounts for conditional uncertainties in biological connections in the given context and gives more nuanced conclusions. The prior study<sup>21</sup> focused primarily on gathering the literature base and developing the Bayes factor to conduct univariate linkage analysis between terms; here, the Bayes factor is used as a tool to consider multivariate relationships and in a high-dimensional Bayesian network, leading to more nuanced and meaningful results.

### 4 Materials and Methods

The proposed framework constructs and analyzes networks based on knowledge embedded in ontological annotations of descriptions of experimental data and abstracts of published literature. After obtaining an annotated knowledge database, the framework comprises three main stages: (1) defining the query, (2) constructing a Bayesian graph based on that query, and (3) using that graph to perform probabilistic inference.

#### 4.1 The Annotated Knowledge Base

**Table 1:** *Sources of records in the knowledge base.* Records were compiled in 2009. For all sources except PubMed, every existing record was included in the knowledge base. For efficiency, 100,000 PubMed records were randomly selected from the 16,000,000 existing at the time. A B-tree index was created on the records for searching.

Source Database	Records	Source Database	Records
Adverse Event Reporting System <sup>14</sup>	774,606	Drug Bank <sup>20</sup>	4774
Array Express <sup>15</sup>	9281	Database of Phenotypes and Genotypes <sup>21</sup>	75,828
BioSiteMaps <sup>16</sup>	1013	Gene Expression Omnibus <sup>22</sup>	15,968
caNanoLab <sup>17</sup>	444	Stanford Microarray Database <sup>23</sup>	16,148
Conserved Domain Databases <sup>18</sup>	34,735	Published articles in PubMed <sup>24</sup>	100,000
Clinical Trials Database <sup>19</sup>	75,828	Drug Bank <sup>20</sup>	4774

We prepared an indexed B-tree for searching the knowledge base that comprised annotated records from eleven corpora

available from NCBO Bioportal in 2009 (Table 1). Then, 220 ontologies were obtained from the NCBO BioPortal; for caching sufficient statistics when searching through the literature, the dictionary of all available ontology concepts (4,153,358 terms) was obtained. More details on preparation of this B-tree and the ontology data are provided in Kshitij et al.<sup>21</sup>

## 4.2 Queries

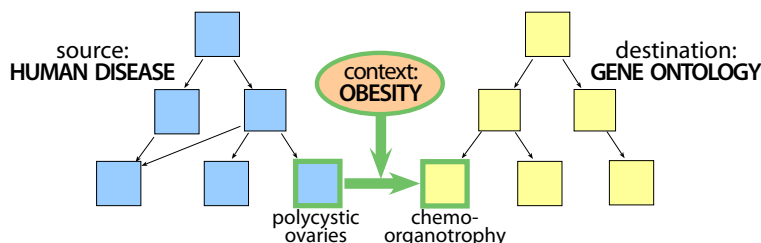
In this work, a *query* consists of a concept of interest (the context under which linkages are identified) and two ontologies (containing the terms between which linkages are drawn). One ontology is designated the source ontology; the other is designated the destination ontology. Users define elements of their queries based on their applications. For example, a researcher interested in obesity-related phenotypes and genes might choose “obesity” for the context and Human Phenotype Ontology and Gene Ontology for the two ontologies.

## 4.3 Constructing the Network

### *Determining the Network Structure*

Based on the query, a tree-augmented naïve (TAN) Bayesian network is constructed, where each node is a random variable that represents the state a specific concept takes in an annotation (either “exists” or “does not exist”). Nodes corresponding to concepts from the source ontology are the parents of nodes corresponding to concepts from the destination ontology. The root node corresponds to the context concept and is a parent of all nodes in the network. The TAN structure is adopted because its requirement that the root node is a parent of every other node parallels the way the context term is present when every ontological connection is identified. The structure is appropriate for context-specific inference.

The time complexity of learning a TAN structure from data using a maximally weighted spanning tree algorithm<sup>25</sup> is  $O(n^2N)$ , where  $n$  is the number of features (the number of concepts in both the source and the destination ontologies), and  $N$  is the number of samples<sup>26</sup>. In this study, the data is a large collection of literature annotated by ontologies. However, the large size of biomedical ontologies renders the use of the original TAN learning structure<sup>25</sup> infeasible. Hence, this framework uses a Bayes factor<sup>23</sup> ( $BF$ ) to identify linkages between any source concept  $S$  and any destination concept  $D$  under a given context concept  $C$ <sup>7</sup>. The higher the Bayes factor, the larger the magnitude of association between two random variables. Therefore, we prune many weak linkages between  $S$  and  $D$  under  $C$  and map only  $S$  and  $D$  concepts that share a mutual  $BF$  greater than a threshold value.



**Figure 1:** *The context-specific ontology-mapping process.* This diagram shows the source term “polycystic ovaries” being mapped to the destination term “chemoorganotrophy” under the context “obesity.” Bold green arrows represent associations between concepts. In general, a source term and a destination term are mapped if and only if they are significantly associated with one another in annotations containing the context concept.

In order to calculate the  $BF$  between any  $S$  and  $D$  considering the context  $C$ , we first create a 2-by-2 contingency table. Each element of this table is determined from the frequencies of co-occurrences of  $S$  and  $D$  in literature that contain  $C$ . Let  $n$  be the number of documents, its subscript ( $S$ ,  $D$ , or  $C$ ) be the type of concept being counted, and the superscript (+ or -) be the state of the concept, where a plus sign (+) signifies “exists” and a minus sign (-) signifies “does not exist.” The contingency table contains  $n^{++}$ ,  $n^{+-}$ ,  $n^{-+}$ , and  $n^{--}$ . Counts are obtained through full-text searches of the knowledge database (Table 1) and are used to calculate  $BF$  using the procedure described in Albert<sup>27</sup>. However,  $BF$  is not calculated for every pair of  $S$  and  $D$ : the hierarchical structure of ontologies allows a more efficient depth-first branch-and-bound algorithm<sup>28</sup> to be used to traverse the two ontologies.

After all significantly co-occurring pairs of  $S$  and  $D$  under the context  $C$  are linked, in accordance to the TAN structure, every concept in the network is linked to the context concept as well. The same destination concept may appear several times, each time linked to a different source concept, because of the TAN requirement that nodes have no more than one non-root parent. The different instances of the same destination concept are not considered as one node because keeping them separate drastically facilitates probabilistic inference.

### Associating Edges with Conditional Probabilities

The network must next be associated with probabilities. For each concept in the net, a table containing the conditional probabilities that each of its states (“exists” or “does not exist”) is true is determined for all combinations of states its parent nodes can take. The conditional probability values are derived from the counts of different combinations of the states of the concept in question and its parents in annotations. For example,  $P(C^+) = n_{C^+} / (n_{C^+} + n_{C^-})$  is one context probability value,  $P(S^+|C^+) = n_{S^+C^+} / (n_{S^+C^+} + n_{S^-C^+})$  is one source probability value, and  $P(D^+|S^+C^+) = n_{D^+S^+C^+} / (n_{D^+S^+C^+} + n_{D^-S^+C^+})$  is one destination probability value. Queries are performed, and counts are collected in the same way as when calculating the Bayes factor to build the network structure. Based on transitive closure of concepts in ontologies, we used the same depth-first branch-and-bound procedure described in Kshittij et al.<sup>21</sup> to prune the ontologies and cache the statistics. This pruning makes the Bayesian network construction efficient enough and scalable to the size of biomedical ontologies.

The final product is thus a three-tiered TAN Bayesian network with the context term at the root, source ontology terms as intermediates, and destination ontology terms as the leaves, related to one another by conditional probabilities based on the frequencies of their co-occurrence in annotations of literature and of experimental data.

The power of these networks comes from Bayesian inference. Because nodes are linked by probabilities, given the prior probability distribution of the root nodes, predictions can be made about the states of any of the other nodes. In this study, Pearl’s message-passing algorithm in trees<sup>25</sup> is implemented so that state information about one or more nodes can propagate along the graph edges and influence the probabilities of the states of other nodes. For example, if certain biological concepts are known to be affected, expressed, or active, the nodes corresponding to those concepts are set to true ( $P(\text{exists}) = 1$ ). The tree is then updated to reflect this new knowledge, and  $P(\text{exists})$  values of all other nodes change accordingly.

#### 4.4 Probabilistic Inference on the Network

##### Identifying Inter-Concept Linkages Using Belief Propagation

The power of these networks lies in Bayesian inference. Because nodes are linked by probabilities, predictions can be made about the states of any of the other nodes given the prior probability distribution of the root nodes. Pearl’s message-passing algorithm<sup>29</sup> is implemented so that state information about one or more nodes can propagate along the graph edges and influence the probabilities of the states of other nodes. For example, if certain biological concepts are known to be affected, expressed, or active, the nodes corresponding to those concepts are set to true ( $P(\text{exists}) = 1$ ). The tree is then updated to reflect this new knowledge, and  $P(\text{exists})$  values of all other nodes change accordingly.

One application of the constructed networks and the proposed inference algorithm is the identification of source or destination concepts in the network that are related to the context  $C$ . To measure the relatedness of a term  $T$  to the context, we associate it with a likelihood ratio  $L$ :

$$L = \frac{P(T \text{ exists} | C \text{ exists})}{P(T \text{ exists} | C \text{ does not exist})} \quad (1)$$

To calculate  $L$ , state of the context node is set to “exists,” and the states of all other nodes are left unknown. Beliefs are then propagated, and  $P(T \text{ exists} | C \text{ exists})$  is found for each node. The context node is then set to “does not exist,” and the other nodes are still left with unknown states. Again, beliefs are propagated, and  $P(T \text{ exists} | C \text{ does not exist})$  is found for each node.  $L$  is the ratio of those two probabilities. It measures how much more likely it is that  $T$  is true when  $C$  is true than when  $C$  is false, not simply how likely it is that the two terms coexist. Hence, a general term such as “disease/disorder” would not score a high  $L$  because there would be little difference in the probability that it exists whether or not the context is true. That is, the terms with the highest  $L$  are most likely to be related specifically to the context and are therefore terms of interest.

A  $p$ -value can also be found for each  $T$ - $C$  link. First, the Bayes factor that  $T$  and  $C$  are associated is determined again in the manner described in Section 4.2, except this time the contingency table contains  $n_{T^+C^+}$ ,  $n_{T^+C^-}$ ,  $n_{T^-C^+}$ , and  $n_{T^-C^-}$ . Using that  $BF$ , an upper bound for the  $p$ -value can be determined as follows<sup>36</sup>, where  $p < 1/e$ :

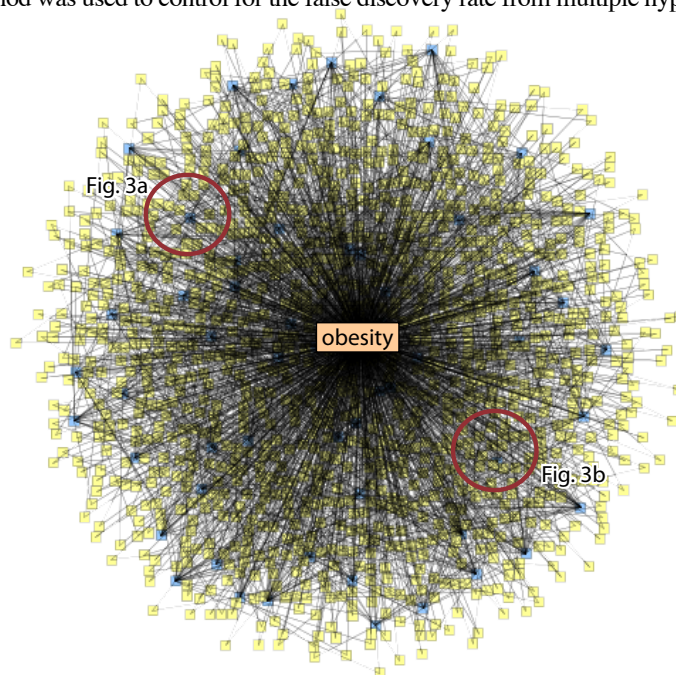
$$-p \ln p < \frac{1}{BF \cdot e}$$

This study examines the terms with the highest  $L$  as the “most related” and then separately associates them with  $p$ -values for ease of understanding. This choice was made because  $L$ , which is based on probabilistic propagation over the network, considers all terms in a high-dimensional joint distribution, whereas using Bayes factor or  $p$ -value as the final mapping is essentially a univariate, deterministic linkage from source to context. Without using  $L$ , the benefits of considering intricate, multivariate biomedical relationships represented in the network would be lost.

### Extending the Network: Gene Inference

Inference using the Bayesian framework is not limited to identification of connections between ontological concepts. For example, the framework can be used to identify the genes most relevant to a queried context. To do so, networks are built with Gene Ontology (GO) as one of the two queried ontologies, and GO concepts in the network are linked with relevant genes based on gene set information from MSigDb<sup>46</sup>. Because links between genes and GO concepts are deterministic, this additional gene level is not actually part of the probabilistic inference framework, and genes do not correspond to network nodes. Therefore, the inference procedure for finding genes relevant to a given context cannot rely on belief propagation. Instead, relatedness of genes to the context are determined based on network structure alone. For each gene, a one-sided Fisher's exact test is used to determine whether there is a significant difference between the proportion of GO terms in the network (which are presumably related to the context) that are associated with that gene and the proportion of GO terms outside the network (which are conversely not strongly related to the context) that are associated with the gene. The Benjamini-Hochberg method was used to control for the false discovery rate from multiple hypotheses<sup>47</sup>.

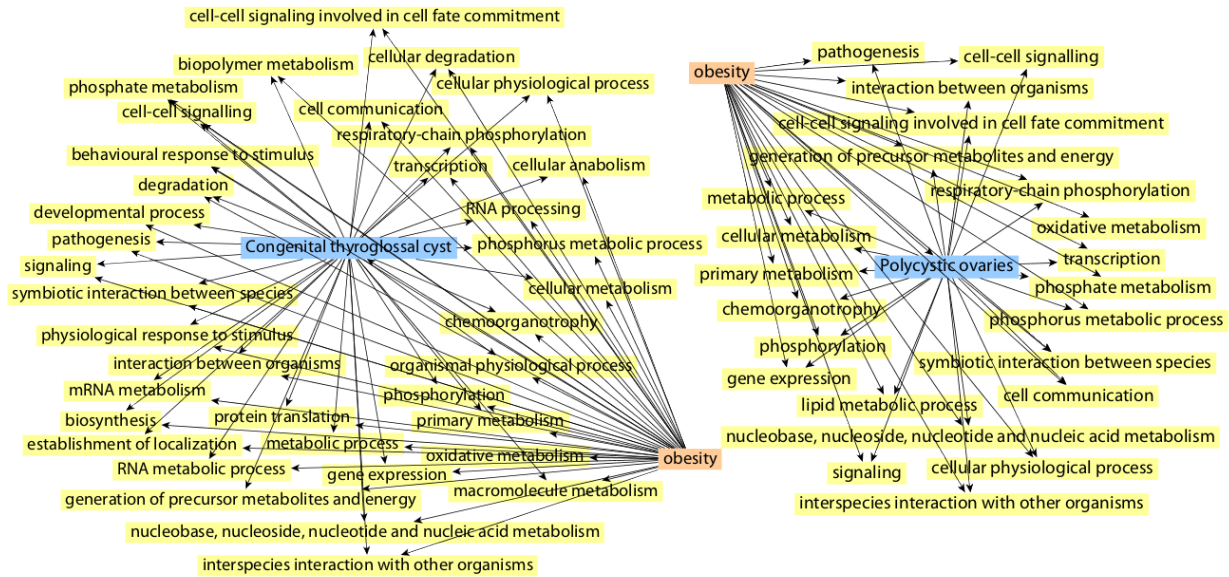
## 5 Results



**Figure 2:** TAN Bayesian network constructed for context “obesity.” Yellow nodes represent GO (destination) terms; light blue nodes represent DOID (source) terms, and the orange node represents the context. Displayed are 4401 out of the 447374 total edges in the network; the full network contains 240 unique nodes from DOID and 8218 unique nodes from GO. Magnified views of the circled subnets are provided in Fig. 3.

In this work, we use the proposed learning network structure and inference procedures to identify diseases and genes related to specific pathologies of interest. The context concept is set to be the pathology, the source ontology is set to be Human Disease (DOID), and the destination ontology is set to be Gene Ontology (GO). Both ontologies are available through NCBO BioPortal<sup>23</sup>. That way, all biomedical relationships represented in the Bayesian network are specific to the context or pathology of interest. Since, as explained in Section 4, inference is done over this context-specific network, all identified relationships are tailored for the context.

Networks were constructed and analyzed for diverse context pathologies, including several cancers, substance abuse disorders, obesity and heart disease, and HIV/AIDS. Similar patterns were observed in the results of all contexts; results from contexts “alcoholism” and “obesity” are discussed further in this paper. The network built using context “obesity” is shown in Figure 2. In the rest of Section 5, italicized body text represents terms or genes identified by the algorithm as associated with the context.



**Figure 3:** Magnified views of circled subnets in Fig. 2. Nodes are color-coded as in Fig. 2. **a)** This subnet is centered about the DOID (source) concept “congenital thyroglossal cyst,” a parent of numerous GO (destination) concepts. **b)** This subnet has the DOID (source) concept “congenital thyroglossal cyst.” True to the TAN structure, the context node “obesity” is a parent of all other nodes as well.

### 5.1 Alcoholism

Disease Term	<i>L</i>	<i>BF</i>	<i>p</i> -value
Drug abuse	340.755	12.949	$4.187 \times 10^{-4}$
Alcohol-related disorder NOS	294.652	12.460	$5.718 \times 10^{-3}$
Substance-related disorder	185.522	11.309	$6.450 \times 10^{-3}$
Disease of environmental origin	74.763	8.707	$8.962 \times 10^{-3}$
Environmentally induced disease	74.763	8.707	$8.962 \times 10^{-3}$
Addiction	68.806	10.934	$6.726 \times 10^{-3}$
Schizophrenia	24.199	5.157	$1.768 \times 10^{-2}$
Alzheimer’s dementia	16.957	3.400	$3.121 \times 10^{-2}$
Organic mental disorder of unknown etiology	6.543	6.341	$1.345 \times 10^{-2}$
Tauopathies	5.114	6.007	$1.445 \times 10^{-2}$

**Table 2.** The ten Disease Ontology (source) terms identified as most strongly linked to the context “alcoholism.”

The disease concepts with the strongest links to the context “alcoholism” using (1) as a measure of link strength are indeed closely biologically related to alcoholism (Table 2). Alcoholism is a *substance-related disorder*, is a form of *addiction*, and would be associated with *alcohol-related disorders NOS*. Alcohol consumption is known to interfere with the nervous system, leading to impaired perception, coordination, memory, and judgment, all possible components of *organic mental disorder of unknown etiology*<sup>38,39</sup>. Psychotic disorders such as *schizophrenia* occur more frequently in alcoholics than in nonalcoholics<sup>40,41</sup>, and alcohol consumption can lead to *tauopathies*, diseases involving aggregation of abnormal tau protein in the brain<sup>42</sup>, such as *Alzheimer’s dementia*<sup>43</sup>. Moreover, environmental factors such as socioeconomic status or education quality play major roles in the development of alcoholism, a *disease of environmental origin* or *environmentally induced disease*<sup>44</sup>, and there exists a high comorbidity between *drug abuse* problems and alcoholism<sup>45</sup>.

The gene inference procedure (Section 4.4) identified many promising genes as significant. For example, a number of genes had already been found by other studies to be associated with the context of alcoholism, including *PTGDS* ( $P < 10^{-15}$ ), the gene with the lowest *p*-value; *MIF* ( $P < 10^{-13}$ ); *BRCA1* ( $P < 10^{-13}$ ); *IL4* ( $P < 10^{-7}$ ); and the three types of peroxisome



proliferator-activated receptor genes (PPARs), *PPARA* ( $P < 10^{-15}$ ), *PPARD* ( $P < 10^{-15}$ ), and *PPARG* ( $P < 10^{-5}$ ). *PTGDS* codes for prostaglandin D2 synthase, which is negatively correlated with alcohol intake<sup>48</sup>. In liver tissues affected by alcoholic liver disease, serum levels of macrophage migration inhibitory factor, coded by *MIF*, are elevated<sup>49</sup>, while alcohol inhibits *IL4*, which controls B-cell proliferation and immunoglobulin class switching<sup>50,51</sup>. Alcohol consumption is associated with heightened incidence of breast cancer, and ethanol down-regulates *BRCA1*, the second most likely gene, of which mutations are closely linked to breast cancer<sup>52,53</sup>. Both *PPARA* and *PPARD* are downregulated by ethanol, *PPARD* agonists alleviate alcohol-induced liver damage, and *PPARG* activation may suppress addictive drinking behaviors<sup>54</sup>. Other significant genes, such as the transcription factor gene *TCF7* ( $P = 6.027 \times 10^{-10}$ ), have not yet been linked in a molecular biological study to ethanol; however, they have been found in other bioinformatics studies to be significantly associated with ethanol or with alcohol withdrawal<sup>55</sup> and therefore are encouraging targets for future biological studies of alcohol dependence.

## 6.2 Obesity

When the same procedure was conducted with the context “obesity,” the algorithm just as successfully identified diseases relevant to the context concept (Table 3). *Obesity, unspecified* is a synonym of the concept itself; *morbid obesity*, defined as weighing 45 kg or more above the ideal weight or having a BMI of at least 40, is a subset of the context<sup>56</sup>. *Polyphagia*, an *eating disorder* characterized by excessive consumption of food, can cause weight gain and lead to obesity. Alcohol intake is another potential cause of obesity<sup>57</sup>, and both alcohol and obesity are associated with fatty liver disease<sup>58</sup> (*alcoholic liver damage, alcohol induced liver disorder*). Obesity increases the risk of *cholelithiasis*, the development of gallstones, especially during the weight loss process<sup>59</sup> and is highly associated with polycystic ovary disease (*ovarian dysfunction, ovarian non-neoplastic disease*), with around 30% of individuals with polycystic ovary disease being obese<sup>60</sup>. There exists a genetic disorder, Ayazi syndrome, characterized by obesity, *choroideremia*, and congenital deafness<sup>61</sup>.

Similar to the case of the context “alcoholism,” the proposed method identified as significant a promising mix of already-corroborated and potentially-related genes for the context “obesity.” For instance, *TGFB1* ( $P < 10^{-7}$ ), a tissue growth factor that regulates proliferation, migration, and differentiation of diverse cells, is linked to abdominal obesity and insulin and glucose imbalance<sup>62</sup>. *PPARD* ( $P < 10^{-7}$ ) activates other genes that direct fatty acid catabolism and thermogenesis; underexpression of *PPARD* results in obesity<sup>63</sup>, while *PPARD* agonists mimic exercise and make promising targets for treatment of metabolic syndromes<sup>63,65</sup>. *UBB* ( $P < 10^{-6}$ ) is one of several genes that codes for ubiquitin, a protein-recycling regulator involved in lipid metabolism and whose levels are inversely associated with BMI<sup>66</sup>. Indeed, mice lacking the *UBB* gene exhibit adult-onset obesity<sup>67</sup>. *CARTPT* ( $P < 10^{-6}$ ) encodes hypothalamic satiety factors<sup>66</sup>, the dysregulation of which may lead to overeating, and *FADS1* ( $P < 10^{-10}$ ), which codes for fatty acid desaturase, is related to lipid metabolism and the plasma triacylglycerol response<sup>69</sup>; both genes easily might relate to obesity. One interesting find was *YWHAH* ( $P < 10^{-9}$ ). Polymorphisms of *YWHAH* are associated with schizophrenia, and antipsychotic drugs<sup>70</sup>, including schizophrenia medications, are known to induce obesity<sup>71</sup>. Perhaps, *YWHAH* is a missing link in knowledge that this method has identified.

Disease Term	<i>L</i>	<i>BF</i>	p-value
Obesity, unspecified	12225.085	630.816	$5.999 \times 10^{-4}$
Polyphagia	11458.037	20.524	$3.102 \times 10^{-3}$
Morbid obesity	5790.621	11.878	$6.067 \times 10^{-3}$
Alcoholic liver damage, unspecified	1416.854	7.704	$1.047 \times 10^{-2}$
Eating disorder, unspecified	163.686	16.905	$3.929 \times 10^{-3}$
Cholelithiasis	123.205	6.733	$1.246 \times 10^{-2}$
Choroideremia	123.205	6.733	$1.246 \times 10^{-2}$
Alcohol induced liver disorder	113.348	6.733	$1.246 \times 10^{-2}$
Ovarian dysfunction	93.180	15.432	$4.392 \times 10^{-3}$
Ovarian non-neoplastic disease	93.180	15.432	$4.392 \times 10^{-3}$

**Table 3.** The ten Disease Ontology (source) terms identified as most strongly linked to the context “obesity.”

## 6 Discussion and Future Work

Our technique can be seen as the first “automatic” probabilistic inference algorithm that uses large biomedical ontologies in conjunction with the vast corpus of existing biomedical literature and experimental data to address specific queries. Of the many probabilistic Bayesian frameworks proposed so far, only this one uses context-specific formulae to map concepts and to calculate conditional probabilities and specializes on a context-specific Bayesian network structure for inference. Therefore, inference using this technique is customized for the researcher’s interests than inference using previous methods.



In this particular work, the proposed framework was effectively used to identify disease concepts and genes related to a context pathology of interest using existing knowledge embedded in literature and in ontologies. Identified disease concepts were invariably closely related to the context. Many of the genes the method identified likewise were known to be associated with the context pathology. Of the remaining genes, many had functions that could logically link them to the context concept or had been identified by other bioinformatics studies as differentially expressed in individuals exhibiting the context pathology. Such genes are promising and interesting because they may constitute new links that augment existing knowledge. The disease concepts and genes identified here may seem to be new information for a researcher with a specific query but no prior information.

All inferences are drawn from the annotated knowledge base that the framework uses as data. Therefore, it is critical that the annotation methods and the selection of data and literature included are comprehensive and representative. We must assume that our knowledge base satisfies the previous condition. Nonetheless, this work advances our ability to generate inferences from such bases. Because literature, experimental data, and ontologies continually evolve, the database must be up-to-date to comprehensively use the prior biomedical knowledge available. We intend to fully automate the data-preparation process in the future and integrate it with the inference framework presented here.

The proposed algorithm can be enhanced to improve its utility as an inferential tool. In this work, a query consists of a context concept of interest and two ontologies of terms from which connections are drawn. In future studies, we intend to extend the algorithm to be able to handle more complex queries. Additionally, future work can examine the predictive power of the framework in identifying drug-disease, drug-pathway, and pathway-disease relationships.

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